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Please find below and/or attached an Office communication concerning this application or proceeding.

	Application No.	Applicant(s)			
		Applicant(s)			
Office Action Summary	09/211,691	GILBERT ET AL.			
Onice Action Summary	Examiner	Art Unit			
· ·	Manjunath N. Rao, Ph.D.	1652			
The MAILING DATE of this communication appeared for Reply	ears on the cover sheet with the	e correspondence address			
A SHORTENED STATUTORY PERIOD FOR REPLY THE MAILING DATE OF THIS COMMUNICATION. - Extensions of time may be available under the provisions of 37 CFR 1.13 after SIX (6) MONTHS from the mailing date of this communication. - If the period for reply specified above is less than thirty (30) days, a reply - If NO period for reply is specified above, the maximum statutory period w - Failure to reply within the set or extended period for reply will, by statute, Any reply received by the Office later than three months after the mailing earned patent term adjustment. See 37 CFR 1.704(b).	within the statutory minimum of thirty (30) ill apply and will expire SIX (6) MONTHS from the application to become ABANDO	timely filed days will be considered timely. om the mailing date of this communication. NED (35 U.S.C. § 133).			
Status		:			
1)⊠ Responsive to communication(s) filed on <u>03 Ma</u>	arch 2004.				
2a)⊠ This action is FINAL . 2b)☐ This	action is non-final.				
3) Since this application is in condition for allowance except for formal matters, prosecution as to the merits is					
closed in accordance with the practice under Ex parte Quayle, 1935 C.D. 11, 453 O.G. 213.					
Disposition of Claims					
4) Claim(s) 37-48 is/are pending in the application 4a) Of the above claim(s) is/are withdraw 5) Claim(s) is/are allowed. 6) Claim(s) 37-48 is/are rejected. 7) Claim(s) is/are objected to. 8) Claim(s) are subject to restriction and/or Application Papers 9) The specification is objected to by the Examiner 10) The drawing(s) filed on is/are: a) acceed applicant may not request that any objection to the or	vn from consideration. election requirement. epted or b) □ objected to by the drawing(s) be held in abeyance.	See 37 CFR 1.85(a).			
Replacement drawing sheet(s) including the correction 11) The oath or declaration is objected to by the Example 11.					
Priority under 35 U.S.C. § 119					
 12) Acknowledgment is made of a claim for foreign a) All b) Some * c) None of: 1. Certified copies of the priority documents 2. Certified copies of the priority documents 3. Copies of the certified copies of the priori application from the International Bureau * See the attached detailed Office action for a list of 	have been received. have been received in Applicative documents have been received in Applicative documents have been received.	ation No ived in this National Stage			
Attachment(s) 1) Notice of References Cited (PTO-892) 2) Notice of Draftsperson's Patent Drawing Review (PTO-948) 3) Information Disclosure Statement(s) (PTO-1449 or PTO/SB/08) Paper No(s)/Mail Date	4) Interview Summa Paper No(s)/Mail 5) Notice of Informa 6) Other:				

Art Unit: 1652

DETAILED ACTION

Claims 37-48 are currently pending and are present for examination.

Applicants' amendments, arguments and declaration under 37 C.F.R. 91.132 filed on 3-1-04, have been fully considered and are deemed to be persuasive to overcome the rejections previously applied. Rejections and/or objections not reiterated from previous office actions are hereby withdrawn. Specifically, Examiner has withdrawn the rejection of claims 1-3, 5-12, 23-27, 33-35, under 35 U.S.C. 103(a) as being unpatentable over Bulow et al. (TIBtech, 1991, Vol. 9:226-231), Defrees et al. (WO 96/32491), and the common knowledge in the art of molecular biology provided by Sambrook et al. (Molecular Cloning, A Laboratory Manual, 2nd Ed, ColdSpring Harbor Laboratory Press, 1989, pages 7.37-7.52) in view of cancellation of above claims. Examiner has withdrawn the provisional rejection of claims 1-3, 5-12, 23-27, 33-35, under 35 U.S.C. 101 as claiming the same invention as that of copending Application No. 10/317,773 (claims1-3, 5-6, 8-12, 23-27, 33-35) in view of cancellation of said claims in the instant application. Examiner has withdrawn the rejection of claims under 35 U.S.C. 112, Ist paragraph as lacking written description in view of the persuasive arguments and the declaration filed by the applicants.

Claim Rejections - 35 USC § 103

The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:

(a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

Claims 37-48 rejected under 35 U.S.C. 103(a) as being unpatentable over Bulow et al. (TIBtech, 1991, Vol. 9:226-231), Defrees et al. (WO 96/32491), Gilbert(a) et al. (Eur. J.

Art Unit: 1652

Biochem., 1997, Vol. 187:187-194) and Gilbert(b) et al. (Biotech. Lett., 1997, Vol. 19(5):417-420) and the common knowledge in the art of molecular biology provided by Sambrook et al. (Molecular Cloning, A Laboratory Manual, 2nd Ed, ColdSpring Harbor Laboratory Press, 1989, pages 7.37-7.52), . Claims 37-48 are drawn to an polynucleotide encoding a fusion polypeptide comprising a bacterial, i.e., a Neisserial, α 2,3-sialyltransferase and CMP-Neu5Ac synthetase, comprising a signal sequence and a molecular tag wherein the two polypeptides are linked through a peptide linker, an expression vector comprising said polynucleotide, a host cell expressing said vector, a method of making said polypeptide by growing the host cells and followed by purification of the fusion polypeptide and permeabilizing the host cell.

Bulow et al. teach the value of artificial bi-functional enzyme as well as multienzyme systems obtained by gene fusion. The reference teaches that preparation of bi and multifunctional enzymes by gene fusion has a great potential in enzyme technology as they facilitate easy purification and exhibit favorable enzyme kinetics. The reference also teaches that selective enzymes can be made as fusion enzymes and used in biochemical analysis, enzyme process technology and metabolic engineering. However the reference does not teach specifically the making of a fusion polynucleotide encoding a fusion protein comprising a sialyltransferase and sialic acid synthetase.

Defrees et al. teach the enzymatic synthesis of glycosidic compounds such as sialic acid compounds using individual enzymes such as a glycosyltransferase, a sialyltransferase and a CMP-NeuAc synthetase for generation of sialic acid and CTP (see entire document and specifically claims 1-10). The reference teaches in detail the other requirements for the enzymatic synthesis of carbohydrate compounds involving multiple enzymes. However, while

Art Unit: 1652

the reference teaches the use of each individual enzymes isolated by recombinant or natural sources the reference does not teach the use of gene fusion either for making these multiple enzymes.

Sambrook et al. provide an exhaustive volume of methods that can be used for various gene manipulations including making fusion polynucleotides, introduction of linker sequences and use of tag sequences in recombinant proteins for easy purification. The reference also teaches purification of recombinant proteins using the molecular tags associated with such proteins. Techniques such as permeabilizing cells for easy access of the encoded enzymes to the substrates are also well k known in the art.

Gilbert(a) et al. teach the characterization of a recombinant *Neisseria* α 2,3-sialyltransferase which plays an important role in the transfer of sialic acid from CMP-NANA to acceptor oligosaccharides which in turn plays a role in cell-cell recognition. The reference also teaches that investigation of the enzymology of glycosyltransferase involved in LOS biosynthesis is limited due to the lack of bacterial glycosyltransferase. The reference provides the amino acid sequence of the enzyme from which a cDNA clone can be developed.

Gilbert(b) et al. teach the purification and characterization of the recombinant CMP-sialic acid synthetase (CSA) from *Neisseria* and teach its use coupled with α 2,3-sialyltransferase(ST) to synthesize CMP-sialic acid which is further attached to various biopolymers. The reference teaches that the major application of the CSA is in "coupled reactions" with sialyltransferases to sialylate oligosaccharides using CTP and NANA as substrates instead of CMP-Neu5Ac which is relatively unstable and expensive. The reference teaches the use of CSA and ST in a coupled reaction to sialylate FCHASE-lactose. The reference concludes that CSA enzyme works

Art Unit: 1652

effectively in a coupled reaction with a ST. The reference also lists several advantages of CSA. However, the reference does not teach the use of fusion polynucleotide comprising encoding sequences of both the above enzymes.

Combining the teachings of the above references it would have been obvious to one of ordinary skill in the art to make a single bi-functional fusion enzyme as taught by Bulow et al. using the CSA and SA taught by Gilbert et al. references. One of ordinary skill in the art would have been motivated to do so in order to develop a one-pot synthesis of FCHASE-lactose. One of ordinary skill in the art would be motivated to make a fusion protein or a host cell comprising a polynucleotide expressing said fusion protein (comprising a Neisseria CSA and ST as taught by Gilbert et al.) because the reference teaches a method involving the use of the two enzymes in the same vessel and that the synthetase works effectively in a coupled reaction with ST. Those skilled in the art would be motivated to permeabilize such host cells using well known methods in the art, so that the enzymes become easily accessible for the acceptor and donor substrates, for direct use of such host cells in sialylating reactions. Furthermore, such a method would obviate the use of expensive and unstable CMP-Neu5Ac from an external source as it is generated in situ and used immediately in the sialylation step. One of ordinary skill in the art would have a reasonable expectation of success since the Gilbert references teach both the enzymes and their compatibility in synthesizing FCHASE-lactose, Bulow et al. teach the increasing use of bi-functional enzymes and Sambrook et al. teach methods for developing fusion protein encoding polynucleotide and methods of making recombinant protein.

Therefore the claimed invention would have been *prima facie* obvious to one of ordinary skill in the art.

Art Unit: 1652

In response to the previous Office action, applicants have traversed the above rejection arguing that none of the cited references disclose all the elements of the claimed invention and also fail to provide motivation for combination of the references to arrive at the claimed invention. Examiner respectfully disagrees with such an argument. First of all Examiner would like to remind the applicants that the above rejection is an obviousness rejection and not based on anticipation and therefore, there is no requirement that each and every reference must teach all the elements of the claimed invention. It is the combination of the teachings of all the references that renders the claimed invention *prima facie* obvious to one of ordinary skill in the art.

Applicants argue that Bulow et al. teaches a number of fusion proteins but do not specifically teach the claimed fusion protein. While that may be so Examiner has used the reference to show that making such fusion proteins was commonly known in the art. Next applicants argue that Defrees et al. teach the method of improving production of sialylated oligosaccharides by using reaction mix comprising unfused alpha-2,3-sialyltransferase and a CMP-NANA synthetase. Here again Examiner used the reference to show that efforts to use the two enzymes together in a single reaction (vessel) to perform sialylation were already existing in the art. With regard to the reference of Gilbert (a) and Gilbert (b), applicants argue that Gilbert(a) fails to provide a motivation for improving the yields of products synthesized by the alpha-2,3,-sialyltransferase and Gilbert(b) et al. do not suggest the combination of alpha-2,3-sialyltransferase with CMP-neu5Ac synthetase and therefore as none of the references either alone or in combination render the claims obvious. Examiner respectfully disagrees with the applicants. As asserted above, while the reference of Gilbert(a) et al. teaches the purification,

Art Unit: 1652

the reference of Gilbert (b) et al. teach the purification and characterization of the recombinant CMP-sialic acid synthetase (CSA) from *Neisseria* and teach its use coupled with α 2,3-sialyltransferase(ST) to synthesize CMP-sialic acid which is further attached to various biopolymers. The reference teaches that the major application of the CSA is in "coupled reactions" with sialyltransferases to sialylate oligosaccharides using CTP and NANA as substrates instead of CMP-Neu5Ac which is relatively unstable and expensive, which by itself is a motivation to use fusion polypeptide comprising the above two enzymes. Applicants argument that references of Gilbert et al. does not render the invention obvious is thus highly misplaced. For all the above reasons Examiner continues to maintain the above rejection.

This application currently names joint inventors. In considering patentability of the claims under 35 U.S.C. 103(a), the examiner presumes that the subject matter of the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation under 37 CFR 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to consider the applicability of 35 U.S.C. 103(c) and potential 35 U.S.C. 102(f) or (g) prior art under 35 U.S.C. 103(a).

Conclusion

None of the claims are allowable.

THIS ACTION IS MADE FINAL. Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE MONTHS from the mailing date of this action. In the event a first reply is filed within TWO MONTHS of the mailing date of this final action and the advisory action is not mailed until after the end of the THREE-MONTH shortened statutory period, then the shortened statutory period will expire on the date the advisory action is mailed, and any extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of the advisory action. In no event, however, will the statutory period for reply expire later than SIX MONTHS from the mailing date of this final action.

Any inquiry concerning this communication or earlier communications from the Examiner should be directed to Manjunath N. Rao, Ph.D. whose telephone number is 571-272-0939. The Examiner can normally be reached on 7.00 a.m. to 3.30 p.m. If attempts to reach the examiner by telephone are unsuccessful, the Examiner's supervisor, Ponnathapura Achutamurthy can be reached on 571-272-0928. The fax phone numbers for the organization where this application or proceeding is assigned is 703-872-9306 for regular communications and for After Final communications. Any inquiry of a general nature or relating to the status of this application or proceeding should be directed to the receptionist whose telephone number is 571-272-1600.

Manjunath N. Rao Ph.D. Patent Examiner, A.U. 1652

5/24/04